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(54) Polyurethane foam laminates.

(57) A structure is disclosed comprising a layer of polyurethane foam bonded to at least one layer of another material, especially a layer of fibreglass matte, a layer of cellulosic material e.g. cardboard, and/or a layer of polyurethane foam. The layers are bonded together with an adhesive of polypropylene that has been grafted with an ethylenically unsaturated dicarboxylic acid or anhydride thereof, the adhesive containing at least 0.03% by weight of said acid or anhydride. The structure exhibits a temperature for heat-fail resistance in shear of at least 90°C. In embodiments, the polyurethane foam is a rigid polyurethane foam, and the polyethylene and polypropylene have been grafted with furnaric acid, maleic acid or maleic anhydride. As an example, the structure is comprised of a layer of fibreglass matte, the layers being bonded together with the adhesive. The structure may be used as so-called headliners in automobiles.

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The present invention relates to laminates of polyurethane foam and especially to laminates of polyurethane foam with fibreglass matte, cellulosic structures e.g. cardboard, and/or polyurethane foam. In particular, the invention relates to such laminates having a temperature for heat-fail resistance in shear, a measurement of the temperature at which detamination of a laminate occurs, of at least 70°C.

Laminates of polyurethane foams with fibreglass matte and/or cardboard are used in a number of enduses, particularly in the form of liners for passenger compartments of automobiles ie. as so-called headliners. Such liners are often multi-layered structures, usually having an outer layer of a fabric material for aesthetic reasons. The traditional adhesive for such structures is a polyurethane adhesive, which has given liners of acceptable structural properties. However, polyurethane adhesives are increasingly in distroour because of environmental and occupational health objections to the use of isocyanate-containing adhesives. Thus, afternate adhesives are required that do comply with environmental and occupational health requirements, as well as meet product specifications for the resultant bonded structure. Moreover, any such adhesive should be economical.

As used herein, a "rigid foam" is defined according to the specifications in ASTM D1566-82 entitled "Definitions of Terms Relating to Rubber".

As used herein, "temperature for heal-fail resistance in shear" is measured according to the procedure of ADM D4498 entitled Heal-Fail Temperature in Shear of Hot-Melf Adhesives, except that the weight used was 300;; use of heavier weights may result in physical failure of the polyurethane foam layer.

It has now been found that polyethylene and polypropylene grafted with an ethylenically unsaturated diocarboxylic acid or anhydride, optionally in the form of grafted polymer blended with ungrafted polymer, may be used in the bonding of polyurethane foam structures, without the use of isocyanates.

Accordingly, the present invention provides a structure comprising:

a layer of polyurethane foam bonded to at least one of a layer of fibreglass matte, a layer of cellulosic material and a layer of polyurethane foam;

said layers being bonded together with an adhesive of polyethylene or polypropylene that has been grafted with an ethylenically unsaturated dicarboxylic acid or anhydride thereof, or derivative thereof, said adhesive containing at least about 0.03% by weight of said acid or anhydride; and said structure exhibiting at temperature for heat-fail resistance in shear of at least 70°C, preferably at least

said structure exhibiting a temperature for neat-rail resistance in shear of at least 70°C, preferably at least 90°C.

In a preferred embodiment of the structure, the polyurethane foam is a rigid polyurethane foam.

In a further embodiment, the polyethylene and polypropylene have been grafted with at least one of fumaric acid, maleic acid, maleic and/did maleic and/did and acrylic acid, or derivatives thereof.

In another embodiment, the structure is comprised of a layer of fibreglass matte, at least one layer of polyurethane foam and a layer of fibreglass matte, the layers being bonded together with said adhesive.

The present invention provides a structure comprising:

a layer of polyurethane foam bonded to at least one of a layer of another material;

said layers being bonded together with an adhesive of polyethylene or polypropylene that has been grafted with an ethylenically unsaturated dicarboxylic acid or anhydride thereof, or derivative thereof, said adhesive containing at least about 0.03% by weight of said acid or anhydride; and

said structure exhibiting a temperature for heat-fail resistance in shear of at least 70°C, preferably at least about 90°C.

The present invention relates to structures of polyurethane foam laminated to other materials, and will be particularly described with reference to laminates to fibregiass matte, cellulosic material and/or polyurethane foam, especially with respect to use of fibregiass matte. The cellulosic material is particularly in the form of a cardboard but other forms of cellulosic material, especially relatively flexible cellulosic material may be used. The surface of the cellulosic material may be treated, provided that any such treatment does not significantly adversely affect the bonding mechanism e.g. polyethylene-coated cardboard may be used. The fibregiass is in the form of a layer of fibregiass matte. It would normally be flexible to the extent that the layer is capable of being bent or shaped to conform to a curved surface e.g. the shape of a headliner in an automobile. The fibregiass may be treated with a finish e.g. a size or primer, as long as the finish does not significantly adversely affect the bonding mechanism. The thickness of the fibregiass matte may be varied over a wide range, although its preferred that the matte be relatively thin, to prevent or reduce the likelihood of detainisation of the matter. In one embodiment, the thickness is about 0.05 cm and the fibregiass matte is a non-woven matte formed from class fibres having a lendth of about 5 cm.

The polyurethane foam may be and preferably is a rigid foam, although it may also be a flexible foam. If the foam is a rigid foam, it preferably possesses a small degree of flexibility, so that it is capable of being conformed to the shape of a curved surface, although an inflexible rigid foam could be formed into the required shape using a thermoforming or other process. The thickness of the polyurethane foam may be varied over a

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wide range, although in one embodiment it is preferred that the foam be relatively thin for convenience and for versatility in the manufacturing process. As an example, in one embodiment the thickness is about 0.5 cm. The pore size of the foam may be varied over a wide range, a preferred range being 0.1-1.0 mm.

The adhesive is a grafted polyethylene or grafted polypropylene. The polymer may be grafted polymer per se or in the form of an un-grafted polymer blended with a grafted polymer. The polymer may be a promopolymer of ethylene or a copolymer of ethylene or a copolymer of ethylene with a minor amount e.g. up to 20% by weight, of at least one higher hydrocarbon alpha-olefin having 3-20, especially 3-10 and particularly 4-10, carbon atoms. Examples of the higher hydrocarbon alpha-olefins include propylene, butnen 1, 4-methy pentinent 1, hexnen-1 and octane-1, and mixtures thereof, or mixtures of two or more of the higher hydrocarbon alpha-olefins. Such polymers may have a density of from about 0.850 g/cm² up to about 0.985 g/cm², especially 0.899 g/cm² up to about 0.950 g/cm² i.e. the polymers include so-called high density polyethylene, low density polyethylene and very low density polyethylene, especially polyethylene having a density of at least 0.940 g/cm². However, the temperature of bonding permissible or achievable in the process of manufacture of the structures may dictate that a polyethylene of lower density, or lower melting point, be used. In addition, the polymers have a melt index, as measured by the procedure of ASTM D1238 (Condition E), in the range of from about 0.1 dg/min to about 200 dg/min, especially 1-80 dg/min and in particular in the range 2-70 dg/min. Alternatively, the polymer may be a polypropylene, especially a homopolymer of propylene or a copolymer of propylene and a minor amount of ethylene. Such polymers are known and are available commercially.

The polymer is grafted with an ethylenically unsaturated dicarboxytic acid or anhydride, or derivatives thereof. Preferred examples of such an acid or anhydride are fumaric acid, malcia anhydride, acrylic acid, carbox acid, carboxide are fumaric acid, malcia anhydride, machide, and acid anhydride, acrylic acid, citraconic acid, citraconic anhydride, itaconic anhydride, endo-bicydo-[2.2.1]-5-heptene-2,3 dicarboxylic acid anhydride, acid achydride, itachydride, itachydride, itachydride, acid anhydride, acid citracydic acid anhydride, and anhydride, anhydride, acid anhydride, acid anhydride, acid anhydride, acid anhydride, and acid anhydride, and acid anhydride, and any acid and separation may be necessary in selecting any particular grafting monomer, depending on the performance requirements of the particular application. Malcic acid and malcic anhydride are especially preferred. The grafted polymer should contain at least about 0.03%, by weight, of the grafted monomer. In preferred embodiments, the grafted polymer contains at least 0.07% and especially at least 0.10% by weight of grafted monomer. Examples of such grafted polymers are available commercially, for instance from Du Pont Canada Inc. under the trade mark Fusabond or may be manufactured by processes known in the art. An example of the latter is the process described in U.S. Patent 4 612 155 of C.S. Wong and R.A. Zelonka, which issued 1986 September 16.

The adhesive may contain additives provided that such additives do not detract from the performance of the adhesive. A variety of materials are commonly used and known in the adhesive art, such as, for example, antioxidants, surface modifiers, stabilizers, fillers, extenders, waxes, foaming agents, pigments, anti-static acents, and crosslinking acents.

Antioxidants generally protect the stability of the adhesive when subjected to heat, or during long term storage in bulk form. Suitable antioxidants include typical hindered phenols such as for example, butylated hydroxytolunes (BHT), ETHYL® 330 and IRGANOX® 1010, believed to be neopentanetetrayl 3,5-di-tert-butyl-4-hydroxy cinnamate, as well as phosphites.

Surface modifiers, such as for example, amide slip agents, such as stearamide, and silica can be used to advantage in adhesive compositions of this invention.

Certain additives customarily employed in adhesive formulations may interfere with the bonding ability of compositions of this invention. The practioner of ordinary skill will be able to determine the negative effect of a potential additive with minimum experimentation.

The structure of the invention is in the form of a laminate of a layer of polyurethane foam bonded or laminated to a layer of fibreglass matte and/or a layer of polyurethane foam. That structure is bonded together by the adhesive described above. In embodiments, the structure is a multi-layered structure. For instance, the structure could be comprised of layers of, in sequence, fibreglass matte, polyurethane foam and fibreglass matte, or a layer of fibreglass matte, for example, the structures could be:

	glass matte adhesive	
 <	polyurethane	foam
	adhesive glass matte	
 <	glass matte	
 <	adhesive polyurethane	form
 <	adhesive	
 <	polyurethane adhesive	
 <	polyurethane adhesive	foam
	glass satte	

In the preferred end-use, the individual leyers are relatively thin e.g. in the range of 0.2-1.0 cm for the polyurethane foam and in the range of 0.02-0.1 cm for fibreglass matte. However, the thickness of the layers is primarily determined by the intended end-use.

The structure may be menufectured by laminating the layers in a press under heat and pressure. For instance, the various layers may be formed into a sandwich of the required construction and then inserted into the press. The temperature and period of time in the press will depend on, for example, the particular construction and thickness thereof as well as the particular grafted polyethylene or polypropylene selected for the adhesive layer. Examples of temperatures and pressures are given in the examples below. Alternatively, the adhesive may be applied in powder form, as a non-woven sheet or by using a porous coat system e.g. a coating system that applies a libre or web e.g. a number of random strands, of adhesive oroto a substrate and essentially forms a non-woven arbeits layer. The adhesive could also be the outer layer of a multi-layer laminate.

In one embodiment of processes for the manufacture of headliners, steam is injected into the mould during the bonding process at a temperature sufficient to melt the adhesive, with the entire headliner panel being under pressure, open-celled foams permit the escape of any steam used during the bonding process. The foam may be formed by the reaction of a polysocyanate with a polyol using a foaming agent that is a gaseous hydrocarbon or flourocarbon, or flother inertigas e.g., nitrogen. Themoforming-type bonding processes, a.g., with the adhesive pre-applied to one of the layers or components of the laminate or inserted separately into the process, may also be used.

The taminated structure is preferably sufficiently flexible so as to be moulded to the shape of a mould, on the structure of the structure of

The structures of the invention have a temperature of heat-fall resistance in shear of at least 70°C and preferably at least 90°C. Such temperatures are important in the automotive industry, for example, because headliners tend to get hot when the automobile is sitting in the sun and acceptable temperature preformance is required.

Although the present invention has been described herein with particular reference to the bonding of glass matte to rigid polyurethane foam, the invention may also be used in the bonding of rigid polyurethane foam, plass matte to flexible polyurethane foam, and other combinations of rigid polyurethane foam, flexible polyurethane foam, flexible polyurethane foam and glass matte, bonding to cellulosic materials, and including the bonding of like and of dissimilar materials.

The present invention is illustrated by the following examples.

Example I

A laminate of fibreglass matte and a polyurethane foam was formed, with the layers being bonded together using a film of adhesive. The fibreglass was a non-woven matte having a thickness of approximately 0.5 mm that had been formed from glass fibres having a length of about 5 cm. The polyurethane foam was a polyether polyurethane identified by the trade mark "Foamex", with a density of about 0.025 g/cm², a thickness of 0.5 cm and pore sizes that were predominantly in the range of 0.25-0.5 mm. The adhesive film was in sheet form, with a thickness of 0 18 mm.

The adhesive was formed from a polymer, and optionally contained a grafted polymer, the grafted polymer, when used, was used as a blend with the indicated polymer in an amount of 10% by weight. The polymer was a polyethylene, the polymers with densities of about 0.95 being homopolymers of ethylene and the polymers with lower densities being copolymers of ethylene and butene-1. The grafted polymer was a high density linear ethylene homopolymer having a density of 0.95 g/cm³, a melt index in the range of 2.0 dg/min and which had been grafted with 1% by weight of maleic anhydride.

The structure was laminated by heating in a press at a temperature of 140°C for 30 seconds under a pressure of 103 kPa.

The laminate obtained was tested using the procedure of ASTM D4488, except that because of the lack of strength of the polyurethane foam under test conditions, the weight used was 300g. One part of the laminate was clamped and the other part, with weight attached, was allowed to hang freely. The sample was located in a temperature controlled oven. The test involved increasing the temperature of the oven until the adhesive bond failed.

Further details and the results obtained are given in Table I.

TABLE I

	Run No.	Polymer Density	Melt Index	Contains Grafted Polymer	Fail Temp. (°C)
20	1	0.895	12	no	82.0
	2	0.930	73	no	99.4
	3	0.959	65	no	86.7
25	1	}			
	4	0.895	12	yes	120
	5	0.930	73	yes	110
30	6	0.959	65	yes	>120

The results show that the presence of maleicanhydride grafted polyolefin greatly improves the temperature performance, as compared to the un-grafted polyolefin composition.

Example II

The procedure of Example I was repeated, bonding a rigid polyurethane foam to rigid polyurethane foam using the maleic-anhydride grafted polyethylenes of Example I. The results obtained are given in Table II.

TABLE II

Run No.	Polymer Density	Melt Index	Contains Grafted Polymer	Fail Temp. (°C)	
7	0.895	12	yes	129.0	
8	0.930	73	yes	130.7	
9	0.959	65	yes	137.7	

The results show that the grafted polyolefin adhesive blend provided outstanding heat resistance.

Example III

Alaminate of fibreglass matte and polyurethane foam was formed, with the layers being bonded together using a film of adhesive, following the procedure of Example I.

The laminates were subjected to a creep test, using samples as prepared in Example I for the shear adhesion fail test of ASTM D4498. The sets sample was placed in a temperature-controlled oven using a 100g weight; one part of the laminate was clamped and the other part, with weight statched, was allowed to hang freely. The samples were kept at a controlled temperature for one month, after which the temperature was raised by 10°C and the procedure repeated. The temperature at which the adhesive bond failed was recorded. Further details and the results obtained are given in Table III.

TABLE III

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	Run No.	Adhesive	Creep Failure Temp (°C)	SAFT*
			remp (C)	(-0)
10	10	λ	140	138
	11	В	140	137
	12	c	>150	149
	13	D	>150	130
	14	E	140	119
15	15	F	110	91
	16	G	110	88
	* SAFT = Heat	Pail Temperature	e in Shear	
	Adhesive A =	85.65% ungrafte	d high density	
20		polyethylene; 1	4% grafted high	density
		polyethylene (0		
		3 dg/min, 1.0%	by weight of ma	leic
		anhydride); 0.3	5 neopentanete	trav) 3.5-
		di-tert-butyl-4	-hydroxy cinnam	ate
25		antioxidant, fi	nal composition	having a
2.5		melt index of 4	.3 dg/min and 0	.14% by
		weight of malei	c anhydride	
	Adhesive $B =$	83.2% ungrafted	high density	
		polyethylene; 1	6.7% grafted hi	qh density
30			-	_
30				
			.9585 g/cm³, me	
			by weight of ma	
			* neopentanetet	
35			-hydroxy cinnam	
			nal composition	
		melt index of 4 weight of malei		.1/# by
	Adhesive C =	77% by weight o		. 200
	VOTTERTAE C 2	weight of linea	or bordbrobatene	; 204 by
40		polyethylene an	d 29 ber majest	of modes
		polypropylene,	the finel compo	or granted
	•	having a melt i		
		containing 0.10		
		anhydride	,	
45	Adhesive D =	90% high densit	v polvethylene	(0.959
		g/cm3 and melt	index of 65 da/z	nin: 10%
		grafted high de	nsity polyethyl	ene
		(0.9583 q/cm3, 1	melt index of 3	dg/min and
		containing 1.0%		
50		anhydride		
	Adhesive $E \approx$	80% high densit	y polyethylene	(0.959
		g/cm3, melt inde	ex of 65 dg/min)	; 20%
		grafted high de	nsity polyethyl	ene
		(0.9585 g/cm3, n	melt index of 3	dg/min and
55		containing 1.0%		
		anhydride)		
	Adhesive F =	zinc ionomer, m	elt index of 5.	2 dq/min
	Adhesive G =	zinc ionom r. m	elt index of 0.	7 dg/min

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The samples of the adhesives of the invention gave substantially better creep results than the samples based on ionomers. The grafted adhesives gave better shear adhesion fall temperatures than the related ungrafted adhesives.

Claims

- 1. A structure comprising:
- a layer of polyurethane foam bonded to at least one of a layer of another material; said layers being bonded together with an adhesive of polyethylene or polypropylene that has been grafted with an ethylenically unsaturated dicarboxylic acid or anhydride thereof, or derivative thereof, said adhesive containing at least about 0.03% by weight of said acid or anhydride; and
- said structure exhibiting a temperature for heat-fail resistance in shear of at least 70°C.
- 15 2. A structure comprising:
 - a layer of polyurethane foam bonded to at least one of a layer of fibreglass matte, a layer of cellulosic material and a layer of polyurethane foam;
 - said layers being bonded together with an adhes/we of polyethylene or polypropylene that has been grafted with an ethylenically unsaturated dicarboxylic acid or anhydride thereof, or derivative thereof, said adhesive containing at least about 0.03% by weight of said acid or anhydride; and
- said structure exhibiting a temperature for heat-fail resistance in shear of at least 70°C.
 - The structure of Claim 1 or Claim 2 in which the temperature for heat-fail resistance in shear is at least 90°C.
 - 4. The structure of any one of Claims 1-3 in which the polyethylene and polypropylene have been grafted with at least one of fumaric acid, maleic acid, maleic anhydride, maleimide, nadic anhydride, acrylic acid, citraconic aintydride, inconic anhydride, incho-bicydo-[22-1]-5-heptene-2,3 dicarboxylic acid, endo-bicydo-[22-1]-5-heptene-2,3 dicarboxylic acid anhydride, cis-4-cyclohexens 1,2 dicarboxylic acid, cis-4-cyclohexens 1,2 dicarboxylic acid, anhydride, and derivatives thereof.
 - The structure of any one of Claims 1-4 in which the structure is comprised of layers of glass matte and polyurethane foam.
- The structure of any one of Claims 1-5 in which the structure is comprised of a layer of fibreglass matte, at least one layer of polyurethane foam and a layer of fibreglass matte, the layers being bonded together with eath of wheelve
 - 7. The structure of any one of Claims 1-6 in which the adhesive is grafted polyethylene.
- 8. The structure of any one of Claims 1-6 in which the adhesive is grafted polypropylene.
 - 9. The structure of any one of Claims 1-7 in which the polyethylene is high density polyethylene.
 - 10. The structure of any one of Claims 1-7 in which the polyethylene is low density or very low density polyethylene.
 - The structure of any one of Claims 1-10 in which the adhesive contains at least 0.7% by weight of grafted acid or anhydride.
- 12. The structure of any one of Claims 1-11 in which the layers of polyurethane foam have a thickness of 0.2-1.0 cm and the fibreolass matte has a thickness of 0.02-0.1 cm.
 - 13. The structure of any one of Claims 1-12 in which the adhesive is formed form a blend of grafted and ungrafted polyethylene or polypropylene.
- 14. The structure of any one of Claims 1-13 in which each layer has sufficient flexibility to be able to be bent or shaped to conform to a curved surface.
 - 15. The structure of any one of Claims 1-13 in which the polyurethane foam is a rigid polyurethane foam.



European Patent Office

EUROPEAN SEARCH REPORT

Application Number EP 94 30 2612

1	DOCUMENTS CONS	DERED TO BE RELEVAN	Г	
Category	Citation of document with i	ndication, where appropriate,	Relevant to chains	CLASSIFICATION OF THE AFFLICATION (bitCLS)
x	US-A-4 645 710 (BA)	TINGER ET AL.)	1,2,4,7, 8,11,15	B32B5/18 B32B7/12
	* column 1, line 56 claims 1-6 * * column 2, line 30	i - column 2, line 8;) - line 61 *	,,,,,,,	
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	* page 6, line 12 - 1,3,8,10 * * page 7, line 8 - * page 10, line 10	line 20 *	11,13	
^	US-A-4 565 733 (FU	II PHOTO FILM CO., LTD.) - line 9; claims 1,2 *	1	
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	* page 1, line 58 - 1,3,6,11 * * page 2, line 40 -	page 2, line 3; claims line 44 *	10,11,10	TECHNICAL FIELDS SEABCHED (SECLS)
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